INTRODUCTION


The morphology and structure of WSSV are well known, and several techniques including histopathology, serology and DNA technology (e.g. PCR and DNA probes) have been developed for its detection. During the past 10 yr, these techniques have been widely used to study modes of WSSV transmission, and several routes of infection have been found. These include not only carrier shrimp larvae, other crustaceans (e.g. crabs) and plankton in and around culture ponds, but also bird feces (Huang et al. 1995a,b, Lo et al. 1996b, Chang et al. 1998, Rajendran et al. 1999, Wang et al. 1999). Despite this knowledge and the implementation of measures to reduce transmission (Wang et al. 1998, 1999), the occurrence of WSSV outbreaks is still very high in mainland China, especially in more northerly regions. This may be due to the fact that some major transmission routes have not yet been discovered.

Huang et al. (1995a,b) reported that some zooplankton, such as copepods, were positive for WSSV by serological detection and might therefore be able to transmit the virus in shrimp ponds. Nowadays, shrimp farmers usually disinfect the pond water before stocking or before addition during the culture process to avoid this problem. In the spring of 2002 at Rushan Shrimp Farm, Shandong, PR China, we disinfected water in 3 culture ponds with high-concentration (100 ppm) bleaching powder before stocking shrimp. In addition, the 3 ponds were enclosed by fences to prevent contact with external water. During the shrimp culture process, no mortality due to WSSV was observed in the 3 ponds. This suggests that the 3 ponds were effectively isolated from outside WSSV sources. However, the occurrence of WSSV outbreaks is still very high in mainland China, especially in more northerly regions. This may be due to the fact that some major transmission routes have not yet been discovered.
prevent crab entry. Despite these measures, we found that new zooplankton that appeared within 20 d post-disinfection (i.e. 1 d before stocking) were positive for WSSV by PCR. Therefore, we hypothesized that some zooplankton resting eggs deposited in the pond sediments may have been a WSSV reservoir with the potential capability of transmitting WSSV through the food web to farmed shrimp. In this study, we examined the possibility that rotifer resting eggs in pond sediments and rotifers hatched from them might be WSSV reservoirs.

MATERIALS AND METHODS

**Sediment collection.** Sediments (1 cm surface layer of pond mud) were collected from 3 ponds in Rushan Shrimp Farm and spooned into plastic bottles to be brought back to the laboratory for analysis.

**Resting egg separation and hatching.** Eggs in the sediments were separated according to the method of Marcus (1989), with slight modification. Briefly, 1 spoonful of mud (approximately 10 ml) was added to a 250 ml beaker containing 200 ml sterilized seawater and sonicated for 30 s at 200 to 400 W using a JY92-II sonifier cell disruptor. The mixture was then filtered through a 52 µm plankton screen and the captured material was resuspended in a solution of refined sugar (sugar:distilled water = 1:1 w/v) and centrifuged for 5 min at 4000 × g. The material remaining in suspension was again filtered through a 52 µm plankton screen and washed thoroughly with seawater before transfer to a dish containing seawater for microscopic observation. Rotifer resting eggs (10) were picked from the dish under a microscope using a capillary tube and placed in a 100 ml beaker with sterilized seawater. Hatching was carried out in an illuminated incubator at 25°C with a 12 h light:12 h darkness regime.

**DNA extraction.** DNA extraction of rotifer resting eggs and hatched rotifers was carried out according to the method of Wang et al. (2000). Materials were stored in SEMP-Tris and extracted by boiling ethanol precipitation. Dried DNA was dissolved with Tris-ethylenediaminetetraacetic acid (TE) buffer.

**Detection methods.** PCR (polymerase chain reaction), DNA probe dot blot hybridization and PCR-inner DNA probe dot blot hybridization were used as detection procedures. Briefly, 2 pairs of primers were designed by Primer Premier 5.0 program (PREMIER Biosoft International, Silicon Valley, USA) based on the whole WSSV DNA sequence (F. Yang et al. 2001). The first primer pair was 5’-CCA AGA CAT ACT AGC GGA TA-3’ and 5’-GAC GAC CCT GAC AGA ATT AC-3’ with a product fragment of 235 bp (256481–256715 bp of AF332093 in GenBank). This pair was used for PCR amplification from DNA templates. The second primer pair was 5’-GGG AAG TGA ATA CGC AGT GA-3’ and 5’-GTT CTA GGG CAA ACA ATG GC-3’, which amplified a 101 bp fragment (256529–256629 bp of AF332093 in GenBank) within the sequence of the first primer pair product. This primer pair was used to prepare a DIG (digoxigenin) labeled DNA probe using DIG Labeling Mix from Roche according to the manufacturer’s instructions. The electrophoresis detection limit of PCR using the first primer pair was approximately 1 pg (about 3000 copies) of WSSV DNA, but this could be improved to 10 fg (about 30 copies) WSSV DNA by dot blot hybridization with the PCR product using the probe described above. In contrast, direct dot blot hybridization could detect only 1 ng (about 3 × 10^6 copies) or more of WSSV DNA (Yan et al. 2004).

**Surface disinfection of resting eggs.** In order to determine whether WSSV was situated within the resting eggs, they were treated for 30 min with Manful King chlorine disinfectant to destroy surface WSSV DNA (B. Yang et al. 2001). They were subsequently washed thoroughly with sterilized seawater before DNA extraction.

RESULTS

**Rotifer resting eggs and hatched rotifers**

Eggs of 5 types, with different shapes and sizes, were separated from Rushan shrimp ponds. At present, only 1 type of rotifer resting egg has been successfully hatched in large quantities; we still do not know what organism produced the 4 other types of unhatched eggs seen. The successfully hatched rotifer resting egg were yellow-brown, 90–110 µm in length and 70–90 µm in width (Fig. 1). This type of egg was rare in

![Fig. 1. Rotifer resting eggs separated from Rushan sediment. Scale bar = 50 µm](image)
June 2002, but abundant in October 2002 and January 2003. Rotifers hatched from these resting eggs (Fig. 2) were identified as *Brachionus urceus*.

**WSSV detection in rotifer resting eggs and rotifers**

Only 1 rotifer resting egg sample containing 3 eggs was WSSV-positive (Lane 3 in Fig. 3) by PCR-electrophoresis. The other 2 resting egg samples (2 eggs and 1 egg) and all rotifer samples (containing 1 to 3 rotifers) were negative. However, PCR-dot blot hybridization with the same PCR products revealed 2 WSSV positive rotifer resting egg samples (A3, B2 in Fig. 4), 4 WSSV-positive rotifer samples (B4, B5, B7, B8 in Fig. 4) and 3 other WSSV-positive egg samples (A4, A7, A10 in Fig. 4). In contrast, 2 samples of seawater used for separating sediment eggs and for washing rotifers (A1, A2 in Fig. 4) were both WSSV-negative. None of the samples gave WSSV-positive results by direct dot blot hybridization without PCR (C1–D9 in Fig. 4).

**Detection of WSSV in resting eggs after surface disinfection**

Testing of rotifer resting egg samples by PCR-dot blot hybridization after Manful King™ disinfection revealed that 8 out of 11 were WSSV-positive (A1–A4, B1–B3, and B5 in Fig. 5); 5 samples of other eggs (C1–C3, C6, and D3 in Fig. 5) were also WSSV-positive by this test. Seawater used to separate and wash the eggs was WSSV-negative (B6 in Fig. 5).

**DISCUSSION**

The fact that WSSV could be detected in rotifer resting eggs and rotifer samples by PCR-dot blot hybridization indicated that WSSV was present in trace amounts and that sensitive methods were necessary to detect it. We tested a total number 16 resting eggs in the surface disinfection test (8 positive samples including 11 eggs and 3 negative samples including 5 eggs). Since some samples were pooled, we do not know whether 1 or more eggs were positive for the pooled samples, but at least 1 of the pooled eggs can be considered positive. Thus, we have a total of 8 positive test results from 16 eggs. Assuming that our samples were a random representation of the whole pond sediments and that the sediments contained in excess of 100,000 rotifer eggs, we used veterinary sampling software (Cameron 2002) to make a rough estimate of the prevalence of WSSV in the eggs. A result of 8 positive tests from a total of 16 would indicate a prevalence of...
at least 72% in the surveyed population. A total of 4 WSSV-positive rotifer samples out of 7 (including a total of 9 rotifers) suggested 4/9 = 44% prevalence. Thus, 72% prevalence in the eggs multiplied by 44% in the hatched rotifers suggested that approximately 32% of the rotifers hatching from eggs in the sampled ponds might be WSSV-positive. Because of the testing method, this may be an underestimate. This is a serious possibility that cannot be ignored.

Pond preparation usually includes a drying period and sometimes soil liming, but specific attention to resting stages of zooplankton present in pond bottoms between crops has been relatively neglected. This is a preliminary report summarizing PCR detection results only, and it has focused primarily on rotifers. Obviously, further work is needed to determine whether rotifers are true carriers of WSSV and capable of transmitting it to shrimp. In addition, follow-up work is needed on other unidentified eggs in the pond sediments that were also WSSV-positive by PCR dot-blot hybridization. This will lead to a better understanding of the risk of WSSV transmission from zooplankton in shrimp pond sediments and the possible development of better prevention methods.

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